

UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER OF PATENTS AND TRADEMARKS Washington, D.C. 20231 www.uspto.gov

APPLICATION NO.	F.	ILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/813,824		03/22/2001	Bert Vogelstein	01107.00112	8421
22907	7590	02/25/2003			
BANNER			EXAMINER		
1001 G STREET N W SUITE 1100 WASHINGTON, DC 20001				CHAKRABARTI, ARUN K	
				ART UNIT	PAPER NUMBER
				1634	
				DATE MAILED: 02/25/2003	

Please find below and/or attached an Office communication concerning this application or proceeding.

Office Action Summary

Application No. 09/813,824

Applicant(s)

Vogelstein

Examiner

Arun Chakrabarti

Art Unit 1634



	The MAILING DATE of this communication appears	on the cover sheet with the correspondence address
	for Reply	
	ORTENED STATUTORY PERIOD FOR REPLY IS SET MAILING DATE OF THIS COMMUNICATION.	TO EXPIRE3 MONTH(S) FROM
	ions of time may be available under the provisions of 37 CFR 1.136 (a). In g date of this communication.	no event, however, may a reply be timely filed after SIX (6) MONTHS from the
- If the p - If NO p - Failure - Any re	period for reply specified above is less than thirty (30) days, a reply within the	and will expire SIX (6) MONTHS from the mailing date of this communication. ne application to become ABANDONED (35 U.S.C. § 133).
Status		
1) 💢	Responsive to communication(s) filed on Nov 8, 20	002
2a) 🗌	This action is FINAL . 2b) 💢 This act	tion is non-final.
3) 🗌	Since this application is in condition for allowance ϵ closed in accordance with the practice under Ex particles.	except for formal matters, prosecution as to the merits is rte Quayle, 1935 C.D. 11; 453 O.G. 213.
Disposi	tion of Claims	
4) 💢	Claim(s) 14-21 and 42	is/are pending in the application.
4	a) Of the above, claim(s)	is/are withdrawn from consideration.
5) 🗆	Claim(s)	is/are allowed.
6) X	Claim(s) 14-17	is/are rejected.
7) 🗶	Claim(s) 18-21 and 42	is/are objected to.
errore.		are subject to restriction and/or election requirement.
	tion Papers	
9) 🗆	The specification is objected to by the Examiner.	
10)	The drawing(s) filed on is/are	a) \square accepted or b) \square objected to by the Examiner.
	Applicant may not request that any objection to the d	rawing(s) be held in abeyance. See 37 CFR 1.85(a).
11)	The proposed drawing correction filed on	is: a) \square approved b) \square disapproved by the Examiner.
	If approved, corrected drawings are required in reply t	to this Office action.
1 2)□	The oath or declaration is objected to by the Exami	ner.
Priority	under 35 U.S.C. §§ 119 and 120	
13)	Acknowledgement is made of a claim for foreign pr	iority under 35 U.S.C. § 119(a)-(d) or (f).
a) 🗀	☐ All b)☐ Some* c)☐ None of:	
•	1. \square Certified copies of the priority documents have	e been received.
2	2. \square Certified copies of the priority documents have	e been received in Application No
	3. Copies of the certified copies of the priority do application from the International Burea se the attached detailed Office action for a list of the	
_	Acknowledgement is made of a claim for domestic	
a) □	1	
	Acknowledgement is made of a claim for domestic	
Attachme		F. C.
	ice of References Cited (PTO-892)	4) Interview Summary (PTO-413) Paper No(s).
2) Not	ice of Draftsperson's Patent Drawing Review (PTO-948)	5) Notice of Informal Patent Application (PTO-152)
3) [] info	rmation Disclosure Statement(s) (PTO-1449) Paper No(s).	6) X Other: Detailed Action

Art Unit: 1634

DETAILED ACTION

Claim Rejections - 35 USC § 112

1. The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

2. Claims 14-17 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention.

The specification discloses compounds having monomer sequence RRRCWWGYYY, nucleotide analogs thereof, single stranded, linear or circular oligonucleotides of human genomic DNA fragments and certain segments of human wild-type p53 protein which corresponds to the p53-specific binding site. Claims 14-17 are directed to encompass any compound that is able to complex specifically with any p53-specific binding site and corresponding sequences from other species, mutated sequences, allelic variants, splice variants, sequences that have a recited degree of identity (similarity, homology), and so forth. None of these sequences meet the written description provision of 35 USC 112, first paragraph. The specification provides insufficient written description to support the genus encompassed by the claim.

Art Unit: 1634

<u>Vas-Cath Inc. v. Mahurkar</u>, 19 USPQ2d 1111, makes clear that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession *of the invention*. The invention is, for purposes of the 'written description' inquiry, whatever is now claimed." (See page 1117.) The specification does not "clearly allow persons of ordinary skill in the art to recognize that [he or she] invented what is claimed." (See <u>Vas-Cath</u> at page 1116.)

With the exception of SEQ ID NO: XXX, the skilled artisan cannot envision the detailed chemical structure of the encompassed polynucleotides and/or proteins, regardless of the complexity or simplicity of the method of isolation. Adequate written description requires more than a mere statement that it is part of the invention and reference to a potential method for isolating it. The nucleic acid itself is required. See Fiers v. Revel, 25 USPQ2d 1601, 1606 (CAFC 1993) and Amgentine.v. Chugai Pharmaceutical Co. Ltd., 18 USPQ2d 1016. In Fiddes v. Baird, 30 USPQ2d 1481, 1483, claims directed to mammalian FGF's were found unpatentable due to lack of written description for the broad class. The specification provided only the bovine sequence.

Finally, <u>University of California v. Eli Lilly and Co.</u>, 43 USPQ2d 1398, 1404, 1405 held that:

...To fulfill the written description requirement, a patent specification must describe an invention and do so in sufficient detail that one skilled in the art can clearly conclude that

"the inventor invented the claimed invention." *Lockwood v. American Airlines, Inc.*, 107 F.3d 1565, 1572, 41 USPQ2d 1961, 1966 (1997); *In re Gosteli*, 872 F.2d 1008, 1012, 10 USPQ2d 1614, 1618 (Fed. Cir. 1989) (" [T]he description must clearly allow persons of ordinary skill in the art to recognize that [the inventor] invented what is claimed."). Thus, an applicant complies with the written description requirement "by describing the invention, with all its claimed limitations, not that which makes it obvious," and by using "such descriptive means as words, structures, figures, diagrams, formulas, etc., that set forth the claimed invention." *Lockwood*, 107 F.3d at 1572, 41 USPQ2d at 1966.

An adequate written description of a DNA, such as the cDNA of the recombinant plasmids and microorganisms of the '525 patent, "requires a precise definition, such as by structure, formula, chemical name, or physical properties," not a mere wish or plan for obtaining the claimed chemical invention. *Fiers v. Revel*, 984 F.2d 1164, 1171, 25 USPQ2d 1601, 1606 (Fed. Cir. 1993). Accordingly, "an adequate written description of a DNA requires more than a mere statement that it is part of the invention and reference to a potential method for isolating it; what is required is a description of the DNA itself." Id. at 1170, 25 USPQ2d at 1606.

The name cDNA is not itself a written description of that DNA; it conveys no distinguishing information concerning its identity. While the example provides a process for obtaining human insulin-encoding cDNA, there is no further information in the patent pertaining to that cDNA's relevant structural or physical characteristics; in other words, it thus does not describe human insulin cDNA. Describing a method of preparing a cDNA or even describing the protein that the cDNA encodes, as the example does, does not necessarily describe the cDNA itself. No sequence information indicating which nucleotides constitute human cDNA appears in the patent, as appears for rat cDNA in Example 5 of the patent. Accordingly, the specification does not provide a written description of the invention of claim 5.

Therefore, only RRRCWWGYYY, nucleotide analogs thereof, single stranded, linear or circular oligonucleotides of human genomic DNA fragments and certain segments of human wild-type p53 protein which corresponds to the p53-specific binding site but not the full breadth of the claim (or none of the sequences and compounds encompassed by the claim) meets the written description provision of 35 USC 112, first paragraph. The species specifically disclosed are not representative of the genus because the genus is highly variant. Applicant is reminded that <u>Vas-Cath</u> makes clear that the written description provision of 35 USC 112 is severable from its enablement provision. (See page 1115.).

Art Unit: 1634

3. Claims 14-17 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention.

Claims 14-17 are rejected under 35 U.S.C. 112, first paragraph, because the specification, does not reasonably provide enablement for the method of providing any compound to any cell which is able to complex specifically with any p53-specific binding site. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the invention commensurate in scope with these claims.

The Court in re Wands, 8 USPQ2d 1400 (CA FC 1988) stated with regard to enablement that

"Factors to be considered in determining whether a disclosure would require undue experimentation have been summarized by the board in Ex parte Forman. They include (1) the quantity of experimentation necessary, (2) the amount of direction or guidance presented, (3) the presence or absence of working examples, (4) the nature of the invention, (5) the state of the prior art, (6) the relative skill of those in the art, (7) the predictability or unpredictability of the art, and (8) the breadth of the claims."

Here, the claims are broadly drawn to method of providing the physiological effect of wild-type p53 protein to a cell by providing any compound to any cell which is able to complex specifically with any p53-specific binding site. However, the specification does not provide

Art Unit: 1634

guidance commensurate in scope with this claim, teaching only one example of binding of human genomic DNA fragments. The specification provides minimal guidance regarding methods for the identification of alternate methodology of any compound (e.g., any proteins or lipids or carbohydrates) in any subjects of any animal species associated with the presence of allelic variant of p53-specific binding sites. There is no working example of any compound or drug trial and its efficacy in any patients of any animal species (including human) associated with the presence of p53-specific binding site. It is highly unpredictable whether or what other compounds would function in the context of highly variant alleles of p53 genes in different human population and different animal species. It is therefore highly unpredictable whether other diagnostic strategies can be identified which meets this specific criteria regarding the method of providing the physiological effect of wild-type p53 protein to a cell by providing any compound to any cell which is able to complex specifically with any p53-specific binding site. Further, drug discovery regiment will be by the trial and error method. This trial and error requirement is borne out because drug discovery is unpredictable and expensive as disclosed by Gryaznov (Column 1, lines 11-15) (U.S. Patent 5,571,903) (November 5, 1996). Further, each compound in any cell of any animal species (including human) associated with the presence of p53 has unpredictable effects on binding and manifestation of physiological effect, and no general method for a priori selection of specific binding with a p53-specific binding site and manifestation of physiological effect is presented. It would require a large amount of experimentation, potentially including the synthesis of billions of chemicals (as only human genome consists of 60,000-100,000 polymorphic or

variable sites), in order to identify additional compounds with the claimed functionality. Given the Wand's factors opposing the full scope of enablement including the limited teaching in the specification, the presence of only one example of human genomic fragments, the teaching of unpredictability in the prior art, the unpredictability of the art, the breadth of the claim, and the large amount of experimentation needed, with only the skill level in the art being neutral towards enablement, it is concluded that undue experimentation is necessary to make and use the invention as broadly claimed.

Allowable Subject Matter

4. Claims 18-21 and 42 are objected to as being dependent upon a rejected base claim, but would be allowable if rewritten in independent form including all of the limitations of the base claim and any intervening claims.

Response to Amendment

5. In response to amendment, all previous 112(second paragraph) rejections and double patenting rejections have been withdrawn. However, new 112(first paragraph) rejections have been included.

Conclusion

6. Any inquiry concerning this communication or earlier communications from

the examiner should be directed to Arun Chakrabarti, Ph.D. whose telephone number is (703) 306-5818. The examiner can normally be reached on 7:00 AM-4:30 PM from Monday to Friday. If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Gary Benzion, can be reached on (703) 308-1119. The fax phone number for this Group is (703) 746-4979. Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the Group analyst Chantae Dessau whose telephone number is (703) 605-1237.

Arun Chakrabarti,

Patent Examiner,

February 20, 2003

How Kr. Chakrabashi
ARUNK. CHAKHADAHII
PATENT EXAMPLES